

TEXAS DEPARTMENT OF HEALTH LICENSING AND ENFORCEMENT DIVISION

BLOODBORNE PATHOGEN CONTROL PROGRAM DEVICE REGISTRATION APPLICATION

Return both the completed application, and **non-refundable** fee made payable to TEXAS DEPARTMENT OF HEALTH, in the envelope provided or mail to: Texas Department of Health, P. O. Box 149200, Austin, Texas 78714-9200. You may visit our website at: www.tdh.state.tx.us/bfds

BBPATHOGEN

BUDGET: ZZ070 FUND: 107

LICENSE #:

MANUFACTURER'S INFORMATION:												
Manufacturer's Name:												
Manufacturer's Mailing Address:												
Manufacturer's Contact Person (Title):												
Manufacturer's Phone #:												
Manufacturer's Fax #:												
Manufacturer's Email Address:												
Manufacturer's Website (URL):												
REGISTRATION	N FEES		one only \$1,500.0		(Non-refu	ŕ	Annual Ren	.1.	- \$1,000.00			
PLEASE NOTE: Registration certificates are not transferable from one device to another or from one device name to another. Any request for transfer of registration due to a change in ownership shall be made in writing to the Texas Department of Health. PRODUCT IDENTIFICATION:												
PRODUCT NAME:												
MODEL NAME AND/OR NUMBER:												
SYRINGE VOLUMES AVAILABLE (if applicable):												
G 1 cc G 3 cc	G 5 cc	G 10 cc	G 20 cc	G 30 cc	G 50 cc	G Insulin	G Tubero	culin	G Other			
NEEDLE GAUGES AVAILABLE (if applicable):												
G 15g G 16g	G 17g	G 18g	G 19g	G 20g	G 21g	G 22g	G 23g	G 25g	G Other			
VERIFICATION: I SWEAR OR AFFIRM THAT ALL OF THE INFORMATION IN THIS APPLICATION IS TRUE AND CORRECT. I FURTHER CERTIFY BY SIGNATURE HEREON; THAT I AM AUTHORIZED TO EXECUTE THIS DOCUMENT ON BEHALF OF THE MANUFACTURER. I UNDERSTAND THAT REGISTRATION OF A NEEDLELESS SYSTEM DEVICE OR SHARPS DEVICE WITH ENGINEERED SHARPS INJURY PROTECTION WITH THE TEXAS DEPARTMENT OF HEALTH, DOES NOT CONSTITUTE AN ENDORSEMENT OR RECOMMENDATION OF THIS DEVICE.												
Signature								Date				
Printed Name & Title												

(Health and Safety Code, Chapter 81, Subchapter H)

Pl			ormation needs to be provided only on i oplication was submitted.	nitial application or if revisions h	ave been made
CC	DMMON NAME/TYPE (Please check only or	ne category an	d one type of device):		
G	Medication delivery devices:	G Vascu	lar access blood drawing devices:	G Surgical/ procedure needles:	
	☐ Disposable syringe injection		Winged, steel-needle IV, butterfly	☐ Type:	
	□ Needleless injection□ Prefilled medication syringe injection		Vacuum tube phlebotomy Arterial blood gas	G Hemodialysis needle set:	
	Other	٥	In-line blood collection Other	☐ Type:	
G	IV Administration:	G Punct	ure/incision administration devices:	G Safety dental syringe:	
	☐ IV needleless administration		Lancet	☐ Type:	
	☐ IV protected needle administration☐ IV catheter (stylet)☐ Other	0	Capillary blood access device Other	G Other	
	Other				<u> </u>
Τŀ	HE DEVICE IS A (check one only):				
G	Needleless System - A device that does not to		nd that is used to withdraw body fluids for any other procedure involving the p		
	risk of an exposure incid or another effective me infusion safety secure	s, accessing a ent by a mech echanism, or i ment device th	vein or artery, or administering medical anism, such as barrier creation, blunting is built into any other type of needle duat effectively reduces the risk of an experience of the risk of	tions or other fluids and that effect, encapsulation, withdrawal, retractive, into a nonneedle sharp, or posure incident.	ctively reduces the action, destruction, r into a nonneedle
	G barrier creation	G bluntii	ng G encapsulation	G withdrawal/retraction	G other
DH	ESCRIBE HOW THE SAFETY FEATURE IS	SACTIVATE	D (if applicable - 300 characters or less)):	
PL	EASE PROVIDE THE FOLLOWING INFO	RMATION W	ITH THE APPLICATION FORM (Che	ck the box to indicate each is encl	losed):
	Brief product description (400 characters or l	ess)			
G G G	Photocopy of labeling submitted to FDA Product marketing or promotional literature Photocopy of original US FDA marketing cle Photocopy of proof of exemption from 510(k) If exempt, provide Code of Federal Regulatio) premarket no	otification (if applicable)	market approval (PMA) submission	on

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